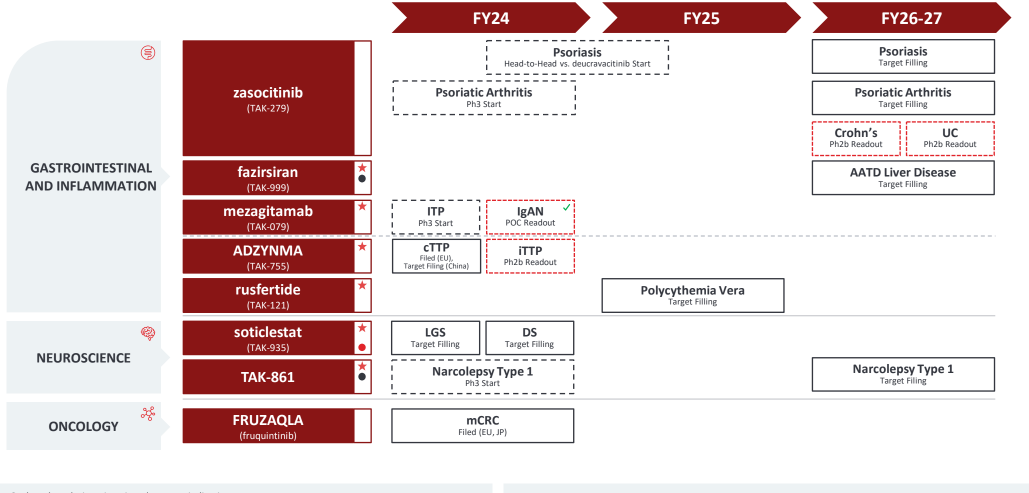
Developing Life Transforming Medicines for Rare and More Prevalent Diseases: Six Late-Stage Programs have the Potential to Generate Significant Value



★ Orphan drug designations in at least one indication

Targeted pivotal study / Phase 3 start

Proof-of-concept/Dose ranging Phase 2 study readout

Target Filing, anticipated year of filing for regulatory approval

Approved

US Breakthrough and/or EU PRIME designations in at least one indication

Japan SAKIGAKE and/or China Breakthrough designations in at least one indication

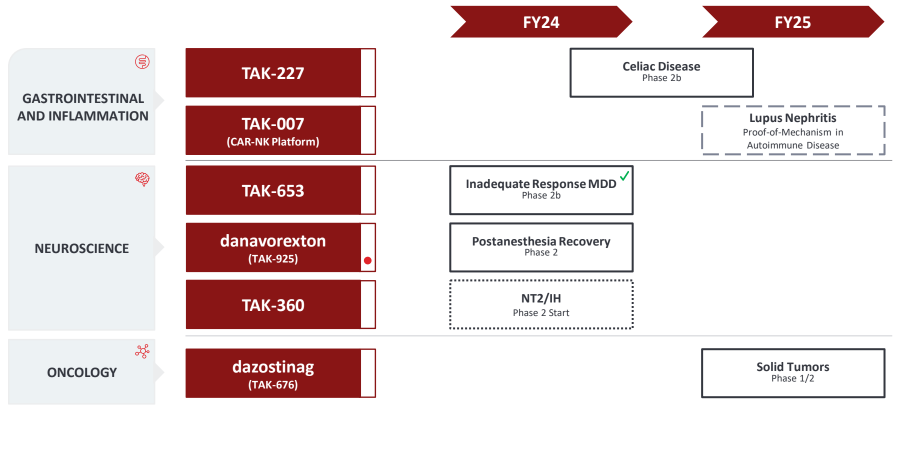
Late-stage program: Program in or expected to be in potential pivotal trial or having achieved proof-of-concept.

Milestone achieved

All timelines are approximate estimates as of May 9th, 2024, are subject to change and are subject to clinical and regulatory success. Table only shows selected R&D milestones and is not comprehensive. For full glossary of abbreviations please refer to appendix.

Impactful Pipeline Milestones for Early to Mid-Stage Programs Address Unmet Patient Needs and Advance Science





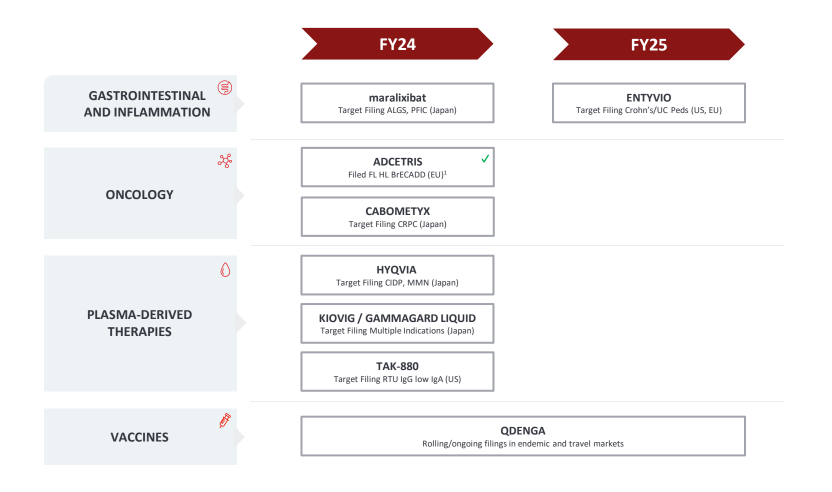
Proof-of-concept (POC): Achieving proof-of-concept means obtaining clinical data sufficient to initiate pivotal trials or late-stage development. A "readout(s)" for a clinical trial occurs when Takeda has (1) received the relevant clinical data, (2) completed any necessary analysis and review of such clinical data, and (3) in instances where it is required or otherwise common convention or practice, consulted with applicable regulatory authorities regarding such clinical data. Where a readout is indicated for a class of related indications (e.g., solid tumors) involving multiple POC clinical trials, such readout occurs upon the earlier of (1) the first achievement of POC in an indication in such class, or (2) the conclusion of all of the POC clinical trials in such class.

[]	Clinical proof-of-mechanism
	Proof-of-concept to inform Go/No-go to pivotal trial
()	Phase 2 Start
•	Japan SAKIGAKE and/or China Breakthrough designations in at least one indication
 Image: A second s	Milestone achieved

All timelines are approximate estimates as of May 9th, 2024, are subject to change and are subject to clinical and regulatory success. Table only shows selected R&D milestones and is not comprehensive. For full glossary of abbreviations please refer to appendix.

Important Near-Term LCM Expansions Represent Significant Growth Opportunities







1. Submission based on data from German Hodgkin Study Group HD21 trial

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Glossary of Abbreviations



Regional Abbreviations: CN: China; EU: Europe; JP: Japan; UK: United Kingdom; U.S.: United States of America American Academy of Dermatology AAD AATD α1-antitrypsin deficiency AATD LD α1-antitrypsin deficiency associated liver disease ACR American College of Rheumatology a disintegrin-like and metalloproteinase with a ADAMTS13 thrombospondin type 1 motifs 13 ADHD attention deficit hyperactivity disorder ALGS Alagille syndrome ALK anaplastic lymphoma kinase ALL acute lymphocytic leukemia AVA Advanced Vial Access BID bis in die, twice a day BLA biologics license application BTD breakthrough therapy designation chimeric antigen receptor natural killer cell CAR NK CHMP Committee for Medicinal Products for Human Use chronic inflammatory demyelinating CIDP polyradiculoneuropathy CML chronic myeloid leukemia CMV cytomegalovirus CPF complex perianal fistulas CRC colorectal cancer CRL complete response letter CRPC castrate-resistant prostate cancer CTCL cutaneous T-cell lymphoma cTTP congenital thrombotic thrombocytopenic purpura DOAC direct oral anti-coagulation DS Dravet syndrome

DSQ	Dysphagia Symptom Questionnaire
EGFR	epidermal growth factor receptor
EMA	European Medicines Agency
EoE	eosinophilic esophagitis
ESS	Epworth Sleepiness Scale
FDA	U.S. Food & Drug Administration
FL	front line
FSI	first subject in
FY	fiscal year
GI	gastrointestinal
GvHD	graft versus host disease
H2H	head-to-head
HAE	hereditary angioedema
HemA	hemophilia A
HL	Hodgkin lymphoma
IARS	International Anesthesia Research Society
IBD	inflammatory bowel disease
IgA	immunoglobulin A
IgAN	immunoglobulin A nephropathy
lgG	immunoglobulin G
IH	idiopathic hypersomnia
IND	investigational new drug
INN	international non-proprietary name
IT	intrathecal
ITP	immune thrombocytopenia
ittp	immune thrombotic thrombocytopenic purpura
IV	intravenous
JAK	Janus kinase
LCM	lifecycle management

LGS	Lennox-Gastaut syndrome
mCRC	metastatic colorectal cancer
mCRPC	metastatic castrate-resistant prostate cancer
MDD	major depressive disorder
MG	myasthenia gravis
MLD	metachromatic leukodystrophy
MM	multiple myeloma
MMN	multifocal motor neuropathy
MSA	multiple system atrophy
MWT	maintenance of wakefulness test
NASH	non-alcoholic steatohepatitis
ND	newly diagnosed
NDA	new drug application
NEJM	New England Journal of Medicine
NK	natural killer
NME	new molecular entity
NMPA	(China's) National Medical Products Administration
NSCLC	non-small cell lung cancer
NT1 or 2	narcolepsy type 1 or 2
PASI	psoriasis area and severity index
PFIC	progressive familial intrahepatic cholestasis
Ph+ ALL	Philadelphia chromosome-positive acute lymphoblastic leukemia
PID	primary immunodeficiency
РК	pharmacokinetics
PMDA	Japan's Pharmaceuticals and Medical Devices Agency
POC	proof of concept
PRIME	Priority medicines scheme by EMA

PTCL-NOS	peripheral T-cell lymphoma not otherwise specified
QD	quaque die, every day
R/R	relapsed/refractory
RTU	ready to use
SC	subcutaneous formulation
SCD	sickle cell disease
SCPCD	severe congenital protein C deficiency
SCT	stem cell transplant
SID	secondary immunodeficiency
SLE	systemic lupus erythematosus
SOC	standard of care
TEAE	treatment emergent adverse event
ткі	tyrosine kinase inhibitor
ттр	thrombotic thrombocytopenic purpura
ТҮК2	tyrosine kinase 2
UC	ulcerative colitis
VEGFR	vascular endothelial growth factor receptors
vWD	von Willebrand disease
WCR	weekly cataplexy rate
ww	Worldwide